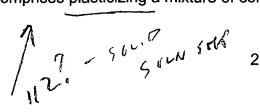
## **CLEAN VERSION OF AMENDMENTS**

## **IN THE CLAIMS**

Cancel claims 5-11.

Add new claims 13-17 as follows:

- phases, wherein one phase forms a matrix which consists essentially of one or more components selected from the group consisting of a homo- or copolymer of N-vinylpyrrolidone, cellulose ether, hydroxyalkylcellulose, celluloseesters and an acrylate- or methacrylate containing polymer and which contains at least one pharmaceutically active ingredient in the form of a solid solution and at least one other phase containing at least one active ingredient is homogeneously incorporated in the form of particles into the matrix phase.
- 14. (new) The composition as claimed in claim 1, wherein said particles are in the form of crystals, pellets, microtablets or granules.
- 15. (new) The composition as claimed in claim 1, wherein a flavoring is present in the matrix phase.
- 16. (new) The composition as claimed in claim 1, which is obtained by incorporating the particles of said other phase into the matrix phase during or after plasticization and shaping the material while still plastic.
- 17. (new) A process for producing the pharmaceutical composition of claim 1 which comprises plasticizing a mixture of components of the matrix phase by energy



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input, incorporating the particles of at least one other phase homogeneously into the matrix phase during or after plasticization, and subsequently shaping the material while still plastic.